AMENDMENTS

In the Claims:

Please add the following claims 184-219:

- 184. The method according to claim 118 wherein the selected samples comprise antibodies that form an antigen-antibody complex with an amino acid sequence of less than about 100 contiguous amino acids encoded by a hepatitis C virus genome.
- 185. The method according to claim 119 wherein the selected samples comprise antibodies that form an antigen-antibody complex with an amino acid sequence of less than about 100 contiguous amino acids encoded by an HCV cDNA insert in the lambda gt-11 library deposited as ATCC deposit No. 40394.
- 186. The method according to claim 123 wherein the selected samples comprise antibodies that form an antigen-antibody complex with an amino acid sequence of less than about 100 contiguous amino acids found in Figure 90.
- 187. The method according to claim 124 wherein the selected samples comprise antibodies that form an antigen-antibody complex with an amino acid sequence of less than about 100 contiguous amino acids found in Figure 14.
- 188. The method according to claim 125 wherein the selected samples comprise antibodies that form an antigen-antibody complex with an amino acid sequence of less than about 100 contiguous amino acids encoded by an HCV cDNA insert in a lambda gt-11 library deposited as ATCC deposit No. 40394.
- 189. A method according to any one of claim 118, 119, 123-125, wherein the selected samples comprise one or more contiguous amino acid sequences selected from the following group:

AA1-AA50; AA1-AA84; AA9-AA177; AA1-AA120; AA35-AA45; AA50-AA100; AA40-AA90; AA65-AA75; AA80-AA90; AA99-AA120; AA95-AA110; AA100-AA150; AA150-AA200; AA200-AA250; AA220-AA240; AA245-AA265; AA250-AA300; AA290-

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AA330; AA290-AA305; AA300-AA-350; AA310-AA330; AA350-AA400; AA405-AA495; AA400-AA450; AA437-AA582; AA450-AA500; AA475-AA495; AA500-AA550; AA511-AA690; AA515-AA550; AA550-AA600; AA550-AA625; AA575-AA605; AA600-AA650; AA600-AA625; AA635-AA665; AA650-AA700; AA645-AA680; AA700-AA750; AA700-AA725; AA725-AA775; AA770-AA790; AA750-AA800; AA800-AA815; AA850-AA875; AA800-AA850; AA920-AA990; AA850-AA900; AA920-AA945; AA940-AA965; AA950-AA1000; AA1000-AA1060; AA1000-AA1050; AA1025-AA1040; AA1075-AA1175-AA1000; AA1000-AA1060; AA1000-AA1050; AA1025-AA1040; AA1075-AA1175; AA1050-AA1200; .AA1070-AA1100; AA1100-AA1140; AA1192-AA1457; AA1195-AA1250; AA1200-AA1225; AA1225-AA1250; AA1250-AA1300; AA1260-AA1310; AA1260-AA1280; AA1266-AA1428; AA1300-AA1350; AA1310-AA1340; AA1345-AA1405; AA1350-AA1400; AA1365-AA1380; AA1380-AA1405; AA1400-AA1450; AA1450-AA1500; AA1475-AA1515; AA1475-AA1500; AA1500-AA1550; AA1515-AA1550; AA1550-AA1600; AA1569-AA1931; AA1570-AA1590; AA1595-AA1610; AA1590-AA1650; AA1610-AA1645; AA1650-AA1690; AA1685-AA1770; AA1689-AA1805; AA1690-AA1720; AA1694-AA1735; AA1720-AA1745; AA1745-AA1770; AA1750-AA1800; AA1775-AA1810; AA1795-AA1850; AA1850-AA1900; AA1900-AA1950; AA1900-AA1920; AA1916-AA2021; AA1920-AA1940; AA1949-AA2124; AA1950-AA2000; AA1950-AA1985; AA2000-AA2050; AA2020-AA2045; AA2045-AA2100; AA2045-AA2070; AA2054-AA2223; AA2070-AA2100; AA2100-AA2150; AA2150-AA2220; AA2200-AA2345; AA2250-AA2330; AA2265-AA2280; AA2280-AA2290; AA2287-AA2385; AA2300-AA2350; AA2350-AA2400; AA2345-AA2415; AA2345-AA2375; AA2348-AA2464; AA2370-AA2410; AA2400-AA2450; AA2400-AA2425; AA2415-AA2450; AA2445-AA2500; AA2371-AA2502; AA2500-AA2550; AA2505-AA2540; AA2550-AA2600; AA2560-AA2580; AA2600-AA2650; AA2620-AA2650; AA2650-AA2700; AA2655-AA2670; AA2670-AA2700; AA2700-AA2750; AA2750-AA2800; AA2755-AA2780; AA2780-AA2830; AA2785-AA2810; AA2796-AA2886;

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Serial Nn. 08/441,355 Docket No. 223002006313 Client Reference PP00063.021 AA2810-AA2825; AA2800-AA2850; AA2850-AA2900; AA2900-AA2950; AA2910-AA2930; and AA2925-AA2950;

wherein the contiguous amino acid sequence is depicted according to the formula $\Lambda A_x - AA_y$, x and y denoting amino acid numbers HCV-1 polyprotein or corresponding regions of other HCV isolates.

- 190. The method according to claim 189 wherein said autibodies are detectable in an ELISA or radioimmunoassay.
- 191. The method according to claim 190 wherein said ELISA or radioimmunoassay employs an antigen comprising said amino acid sequence made by recombinant expression.
 - 192. The method according to claim 190 wherein said biological samples are blood.
 - 193. The method according to claim 191 wherein said biological samples are blood.
 - 194. The method according to claim 190 wherein said biological samples are plasma.
 - 195. The method according to claim 191 wherein said biological samples are plasma.
 - 196. The method according to claim 190 wherein said biological samples are sera.
 - 197. The method according to claim 191 wherein said biological samples are sera.
- 198. The method according to claim 192 wherein the selecting is to identify an HCV positive sample for removal from the supply.
- 199. The method according to claim 193 wherein the selecting is to identify an HCV positive sample for removal from the supply.
- 200. The method according to claim 194 wherein the selecting is to identify an HCV positive sample for removal from the supply.
- 201. The method according to claim 195 wherein the selecting is to identify an HCV positive sample for removal from the supply.
- 202. The method according to claim 196 wherein the selecting is to identify an HCV positive sample for removal from the supply.

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- 203. The method according to claim 197 wherein the selecting is to identify an HCV positive sample for removal from the supply.
- 204. A method according to any one of claim 118, 119, 123-125, wherein the selected samples comprise one or more contiguous amino acid sequences selected from the following group:

AA1-AA84; AA437-AA58?; AA511-AA690; AA9-AA177; AA1192-AA1457;
AA1266-AA1428; AA1694-AA1735; AA1689-AA1805; AA1916-AA2021; AA1949-AA2124;
AA2054-AA2223; AA2200-AA3325; AA2287-AA2385; AA2348-AA2464; AA2371-AA2502;
AA2796-AA2886; AA1569-AA1931,

wherein the contiguous amino acid sequence is depicted according to the formula

AA_x-AA_y, x and y denoting amino acid numbers HCV-1 polyprotein or corresponding regions of other HCV isolates.

- 205. The method according to claim 204 wherein said antibodies are detectable in an ELISA or radioimmunoassay.
- 206. The method according to claim 205 wherein said ELISA or radioimmunoassay employs an antigen comprising said amino acid sequence made by recombinant expression.
 - 207. The method according to claim 205 wherein said biological samples are blood.
 - 208. The method according to claim 206 wherein said biological samples are blood.
 - 209. The method according to claim 205 wherein said biological samples are plasma.
 - 210. The method according to claim 206 wherein said biological samples are plasma.
 - 211. The method according to claim 205 wherein said hiological samples are sera.
 - 212. The method according to claim 206 wherein said biological samples are sera.
- 213. The method according to claim 207 wherein the selecting is to identify an HCV positive sample for removal from the supply.
- 214. The method according to claim 208 wherein the selecting is to identify an HCV positive sample for removal from the supply.

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- 215. The method according to claim 209 wherein the selecting is to identify an HCV positive sample for removal from the supply.
- 216. The method according to claim 210 wherein the selecting is to identify an HCV positive sample for removal from the supply.
- 217. The method according to claim 211 wherein the selecting is to identify an HCV positive sample for removal from the supply.
- 218. The method according to claim 212 wherein the selecting is to identify an HCV positive sample for removal from the supply.
- 219. A method according to any one of claim 118, 119, 123-125, wherein the selected samples comprise one or more contiguous amino acid sequences selected from the following group:

AA1694-AA1735; AA1569-AA1931; AA1192-AA1457; AA1-AA84; and AA9-AA177, wherein the contiguous amino acid sequence is depicted according to the formula AA_x-AA_y, x and y denoting amino acid numbers of HCV-1 polyprotein or corresponding regions of other HCV isolates.

A method of selecting biological samples from a supply of biological samples comprising selecting from said supply those samples that contain a detectable polynucleotide comprising a contiguous sequence of at least 15 nucleotides from either strand of at least one of the hepatitis C virus (HCV) cDNA inserts in a lambda gt-11 cDNA library deposited as ATCC No. 40394.

- 221. The method according to claim 115, wherein the detectable polynucleotide comprises a contiguous sequence of less than about 90 nucleotides fully complementary to either strand of Figure 3.
- 222. The method according to claim 116, wherein the detectable polynucleotide comprises a contiguous sequence of less than about 90 nucleotides fully complementary to either strand of Figure 62A.

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- The method according to claim 117, wherein the detectable polynucleotide comprises a contiguous sequence of less than about 90 nucleotides fully complementary to either strand of Figure 89.
- 224. The method according to claim 118, wherein the selected samples comprise a polymorleotide that hybridizes under stringent conditions to a polymorleotide that comprises a contiguous sequence of less than about 90 nucleotides from the genome of a hepatitis C virus genome or the complement thereof.
- 225. The method according to claim 119, wherein the selected samples comprise a polynucleotide that hybridizes under stringent conditions to a contiguous sequence of less than about 90 nucleotides from either strand of at least one of the hepatitis C virus (HCV) cDNA inserts in a lambda gt-11 cDNA library deposited as ATCC No. 40394.
- 226. The method according to claim 120, wherein the selected samples comprise a polynucleotide that hybridizes under stringent conditions to a contiguous sequence of less than about 90 nucleotides fully complementary to either strand of Figure 89.
- 227. The method according to claim 121, wherein the selected samples comprise a polynucleotide that hybridizes under stringent conditions to a contiguous sequence of less than about 90 nucleotides fully complementary to either strand of Figure 14.
- 228. The method according to claim 115, wherein the detectable polynucleotide comprises a contiguous sequence of less than about 90 nucleotides from either strand of at least one of the hepatitis C virus (HCV) cDNA inserts in a lambda gt-11 cDNA library deposited as ATCC No. 40394.

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